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A cDNA coding for human normal serum albumin a, and a process for production of the albumin.

(iii) A CDNA coding for human hormal serum situmin A; an expression certain comprising the cDNA coding for human expression vertex comprising the cDNA coding for human expression vertex comprising the cDNA coding for human normal serum albumin A; and a process for the production of the human normal serum albumin A; and a process for the production of culturing a host transformed with an expression vector comprising a cDNA coding for the form of a fused with another protein, and obtaining the human normal serum albumin.

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Description

In the drawings:-

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A CDNA CODING FOR HUMAN NORMAL SERUM ALBUMIN A, AND A PROCESS FOR PRODUCTION OF THE ALBUMIN

The present invention relates to a process for the production of human normal serum albumin A by a recombinant DNA technique, and a gene therefor, According to the present invention, a large amount of human normal serum albumin A free of Infection by pathogens such as hepatitis B virus, and AIDS wirus HTLV, can be produced at a low cost

Human serum albumin is a plasma protein synthesized in the liver, and plays an important role in an organism: It serves in the plasma to maintain osmotic pressure; binding various substances such as fatty acids, metal lons such as Cu², M², bile billution, various drugs and water soluble vitamins and the like, to transport same to target organs thereof; and as a source of amino acids provided to tissues. On the basis of such actions, a large amount of human serum albumin is used to treat patients suffering from hemorrhagic shock and hypoalbuminemia generated by a reduced synthesis of albumin due to hepatocirrhosis, or by burns or poshfill;

An amino acid sequence of human normal serum albumin A is known on the basis of an amino acid analysis of natural human serum albumin, and further, CNAs coding for human serum albumin are known. However, amino acid sequences of such polypepides encoded by the known cDNAs are not completely the same as an amino acid sequence of human normal serum albumin A present in most human population. For example, cDNA described by Dugalczyk et al., Proc. Natl. Acad. Sci. USA, 79, 71 - 75 (1982) encodes Gly as the 97th amino acid, although Glu is in human normal serum albumin; cDNA described by Ausmet at., Nucleic Acids Res. 9, 6103 - 6114 (1981) encodes Lys as the 98th amino acid, although Glu is in human normal serum albumin abournin; cDNA described by Mariotit et al., Protides Biol. Fluide Proc. Colloq. 33, 177 - 179 (1985) encodes Tyr as the 92nd amino acid, although As is in human normal serum albumin, and Met as the 381st and 462nd amino acid, although Val is in human normal serum albumin at bath positions; and cDNA described in Japanese unexamined Patent Publication No. 58-150517 encodes Ser as the 98th amino acid, although Cys is in human normal serum albumin acid salven to been determined. A chromosomal DNA sequence coding for human normal serum albumin a disc have to been determined. A chromosomal DNA sequence coding for human normal serum albumin A is described by Minghetti et al., J. Biol. Chem. 251, 674 - 6757 (1986).

Accordingly the above-mentioned cDNAs cannot be used to produce a protein having the same amino acid sequence as human normal serum albumin A.

However, when serum albumin having an amino acid sequence different from that of normal serum albumin is administered to a human, it may exhibit antigenitify and may not exhibit the normal functions of serval albumin, or may have short life time in the blood. Therefore, there is a strong demand for the obtaining of cDNA correctly encoding an amino acid sequence of human normal serum albumin to produce the human normal serum albumin by a recombinant DNA technique.

Accordingly, the present invention provides a cDNA coding for human normal serum albumin A having an amino aoid sequence represented in Figures 3-1 to 3-5.

Moreover, the present invention provides an expression plasmid comprising a cDNA coding for human normal serum albumin A having an amino acid sequence represented in Figures 3-1 to 3-5.

normal serum albumin A having an amino acid sequence represented in Figures 3-1 to 3-5.

The present invention also provides a host transformed with an expression plasmid comprising a cDNA coding for human normal serum albumin A having an amino acid sequence represented in Figures 3-1 to 3-5.

The present invention still further provides a process for the production of human normal serum albumin A comprising the steps of culturing a host transformed with an expression plasmid comprising a CDNA coding for the human normal serum albumin A having a mainine add sequence represented in Figures 3-1 to 3-5, to express the protein alone or in a form of a fused protein with other protein, and obtaining the human normal serum albumin A.

Figure 1 shows restriction enzyme cleavage maps of a cDNA fragment (HSAcDNA) coding for an entire human normal serum albumin A of the present invention, as well as a cDNA fragment (HSA-IA) coding for 3'-terminal side and a cDNA fragment (HSA-II) coding for 05'-terminal side;

Figs. 2-1 to 2-2 show a construction process of various plasmids related to the present Invention; Figs. 3-1 to 3-5 show a nucleotide sequence coding for an entire human normal serum albumin A of the

present invention, and an amino acid sequence corresponding to the nucleotide sequence; Fig. 4 shows a result of an electrophoresis of an expression product representing proteins reacted with

an anti-human serum albumin antibody; and, Fig. 5 shows a nucleotide sequence of three probes used to screen a cDNA library.

A cDNA coding for human normal serum albumin A can be obtained by screening a human cDNA library by a conventional procedura; for example, a human liver library prepared using phage Agr11 as a vector. Probes for screening the cDNA library can be designed on the basis of a known nucleotide sequence of cDNA coding for human serum albumin. Preferably, a combination of three probes coding for an N-terminal region, cannot be certainal region, cannot be certainal region, and C-terminal region of the human serum albumin respectively is used. To obtain DNA correctly coding for an entire human normal serum albumin A, conveniently, different cDNA fragments coding for different parts of human serum albumin A are selected and sequenced, and after confirming that they correctly

encode corresponding parts of the human normal serum albumin, appropriate parts thereof are joined to form an entire DNA. Where it is found that a part of a cDNA fragment does not correctly encode a corresponding part of an amino acid sequence of the human normal serum albumin A, the part of the cDNA not correctly encoding the amino acid sequence is replaced by a cDNA fraction which correctly encodes the amino acid sequence in question, to construct a correct entire cDNA. Where the construction of an entire cDNA from partial cDNA fragments is difficult, the part of DNA not obtained from cDNA can be supplemented by a synthetic double-stranded DNA fragment.

The cDNA coding for human normal serum albumin A by Itself of the present invention can be expressed. Alternatively, the cDNA of the present invention can be joined with other DNA coding for other peptides to express the human normal serum albumin in the form of a fused protein. As a partner for such a fused protein, avirous peptides can be used, and as an example of the partner peptide, a signal peptide of E_ool alkaline phosphatase can be mentioned. Where the human normal serum albumin A is expressed as a fused protein, the signal peptide can be eliminated from the fused protein after the expression, to obtain the desired human

normal serum albumin A.

To express the human normal serum albumin A as a fused protein, cDNA coding for the fused protein is inserted into an expression vector, which is then introduced into a host. As a host for the expression, eukaryotic cells such as animal cells, yeast cells, and bacterial cells can be used, and a vector is selected according to a host selected. An expression plasmid comprises expression control regions including a promoter and Shime-Dalgamo (50) sequence, followed by CDNA coding for the human normal serum albumin.

As the promoter, a trip promoter, lac promoter, \(\lambda\) page promoter such as P₀ or P₀, tufB promoter or mal promoter, or a hybride promoter constructed from said promoters, such as an tac promoter, can be mentioned. As the SD sequence, aDNA sequence corresponding to a sequence in mRNA complementary to the 3'-terminal nucleotide sequence of E_coil 16S RNA is known to be effective for a start of the translation. Atternatively, a completely complementary symbole DNA transpanent can be used as an SD sequence.

Transformation of a host such as E. coll with an expression plasmid can be carried out by a conventional procedure. A transformed host such as E. coll is cultured by a conventional procedure. When E. coll cells are grown to a predetermined cell concentration, it is induced to express the desired gene. The method of induction depends on promoter used, for example, a trp promoter is used and 3-indoleacrylic acid is added to the culture to induce the expression.

Where E.coli is used as a host, the desired protein is intracellularly accumulated. Therefore, to recover the desired protein, the cultured cells are collected, wheely, essuperined in water or a buffer, and then disrupted. Since the desired protein is entained in an insoluble fraction, the insoluble fraction is collected by, for example, centrifugation of titration, and if necessary, weahed. Next, the recovered insoluble fraction is put into a protein solublaring buffer such as a buffer containing sodium dodecyl sulfate and 2-mercaptoethanol to

Next, from the resulting solution containing a fused protein comprising the human normal serum albumin, the profels in excovered and purified by a conventional procedure. The fused protein can be cleaved by, for example, E. coil leader peptidase (signal peptidase I) in vitro, to obtain the desired human normal serum albumin A, by a procedure described by Zwiznaki C, cand Wickner, W, J, Blot. Chem. 255, 7973 (1985).

Examples

The present invention will be further illustrated by, but is by no means limited to, the following examples.

Example 1. Screening of clones containing cDNA coding for human normal serum albumin A A human liver cDNA library constructed using a vector phage \(\frac{\lambda}{gt11} \) commercially available from Clontech, U.S.A. was used to select clones containing a cDNA fragment coding for human normal serum albumin A by plaque hybridization. The Agt11 recombinant phage of the library was infected to E. coli Y1090, which was then plated on an LM agar medium to form 5.5 x 105 transformant plaques. Recombinant DNAs in the plaques were transfered onto membrane filters (Hybond-N; Amersham), and screened using three synthesized oligonucleotide probes labeled with ³²P (specific radioactivity ≥ 10⁷ cpm/µg) by a method of Benton and Davis, Science, 196, 180 - 182 (1977). These three probes are a probe HSA-1 corresponding to a 5'-non-coding region and a 5'- coding region starting 12 base-pairs upstream from ATG start codon and ending at in a codon for 9th amino acid leucine; a probe HSA-2 coding for 248th glycine to 260th leucine; and a probe HSA-3 comprising a 3'-terminal coding region and a 3'-terminal non-coding region starting with a codon for 576th valine and ending 9 nucleotides downstream from the C-terminal leucine codon, all described by Lawn et al., Nucleic Acids Res. 9, 6103 - 6114 (1981). The nucleodide sequence used as probes were on the complementary or negative strand. The nucleotide sequences of these three probes are shown in Fig. 5. These oligonucleotide probes were synthesized by an automatic DNA synthesizer, and labeled using [y-32P] ATP and polynucleotide kinase. Among 200 \(\lambda\gammath{t11}\) clones which gave a positive signal with the probe HSA-2, from 4 clones, DNA was prepared by a method of Blattner et al., Science, 202, 1279-1284 (1978), and digested with EcoRI, and a Southern blot of the digested product was allowed to hybridize with the probe HSA-2 by a method of Southern, J. Mol. Blol. 98, 503 - 517 (1975). DNA fragments having a size of 1.8 Kb, 1.4 Kb, and 1.3 Kb, respectively, were hybridized with the probe HSA-2. Among these, DNA fragments of 1.8 Kb and 1.3 Kb were subcloned in vector pUC19, and these subclones were subjected to colony hybridization using probes HSA-1 and HSA-3, by a method of Grunstein and Hogness, Proc. Natl. Acad. Sci. USA, 72, 3961 - 3965 (1975).

As a result, a clone Agt11 (HSAI-A) which was hybridized with only HSA-3 was obtained. DNA in this clone was digested with various restriction enzymes, and the resulting DNA fragments were inserted into phage vectors M15mP18 and M15m19 RF DNA, and a nucleotide sequence of the DNA was determined by a dideoxy chain termination method of Sanger, F., Nickien, S. and Coulson, A.R. Proc. Natl. Acad. Sci. USA, 74, 5463 - 5467

On the other hand, among the dones which gave a positive signal in plaque hybridization of Agi11 clones using the HSA-2 probe, 20 clones were subjected to plaque hybridization using the HSA-1 probe, and a positive clone Agi11 (HSA-II) was obtained. From this clone, phage DNA was prepared and digested with EcoRI. The digestion product was subjected to Southern hybridization using the HSA-1 probe, and a DNA fragment of 12-SK bedispinated HSA-II was cloud to hybridize with the HSA-1 probe. A nuclectific sequence of this DNA fragment was determined by a dideoxy chain termination method. The HSA-II did not hybridize with

As a result, it was found that the HSA-II lacks a DNA portion coding for the C-terminal portion of human serum albumin, and the HSA-I-A lacks a DNA portion coding for the N-terminal portion of human serum albumin and containing an opal codon TGA as a stop codon in place of the codon TGA coding for 504th series. Restriction enzyme cleavage maps of these DNA tragments are shown in Fig. 1. In these maps, exact positions of restriction enzyme recognizing sites were obtained from a finally determined nucleotide sequence.

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Table 1

Difference between amino acid sequences encoded by human serum albumin cDNA, and by human chromosomal gene, respectively, and amino acid sequence of human normal serum albumin A purifiled from human serum

DNA or Protein (Reference)		Position	of	amino	acid	resi	dues
		92	97	369	381	396	462
Chromosomal DNA	(1)	Ala	Glu	Cys	Val	Glu	Val
cDNA-1	(2)	Ala	Gly	Cys	Val	Glu	Val
cDNA-2	(3)	Ala	Glu	Cys	Val	Lys	Val
cDNA-3	(4)	Thr1)	Glu	Cys	Met	Glu	Met
cDNA-HSA-A	(5)	Ala	Glu	Cys	Val	Glu	Val
Serum Protein-1	(6)	Ala	Glu	Сув	Val	Glu	Val
cDNA-4	(7)	_ 2)	-	Ser	Val	Glu	-

- Underlined amino acids are different from those of human normal serum albumin.
- The amino acid and nucleotide are not described.

Reference

- (1) Minghetti et al., J. Biol. Chem. <u>261</u>, 6747 - 6757 (1986)
- (2) Dugaiczyk et al., Proc. Natl. Acad. Sci. USA 79, 71 - 75 (1982)
- (3) Lawn et al., Nucleic Acids Res. 9, 6103
 6114 (1981)
- (4) Marisitti et al., Protides Biol. Fluids

Proc. Collog., 33, 177 - 179 (1985)

- (5) Present invention
- (6) Takahashi et al., Proc. Natl. Acad. Sci. USA 84. 4413 - 4417 (1987)
- (7) Japanese Unexamined Patent Publication 58-150517

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Example 2. Construction of plasmid pUC.phoA

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A plasmid pUC.phoA containing a synthetic DNA fragment coding for signal peptide of E. coll alkaline phosphatase was constructed as follows.

A DNA fragment having the following nucleotide sequence coding for signal peptide of E. coli alkaline phosphatase was constructed from chemically synthesized oligonucleotide fragments.

	ECOF	RI.								30
25	AA	TTC	ATG	AAA	CAA	AGC	ACT	ATT	GCA	CTG
		G	TAC	TTT	GTT	TCG	TGA	TAA	CGT	GAC
			Met	Lvs	Gln	Ser	Thr	Ile	Ala	Leu
				-						60
30	GCA	CTC	TTA	CCG	TTA	CTG	TTT	ACC	CCT	GTG
	CGT	GAG	TAA	GGC	AAT	GAC	AAA	TGG	GGA	CAC
	Ala	Leu	Leu	Pro	Leu	Leu	Phe	Thr	Pro	Val
			3	NaeI						
35	ACA	AAA	GCC	GGC	G					
00	TGT	TTT	CGG	CCG	C T	TA.	A			
	Thr	Lys	Ala							
				HP	aII	EcoR	I			

The EcoRI sites of both ends of the DNA fragments were provided to insert the fragment into the EcoRI site of a pUC series plasmid, the Hpall after was provided to fuse It with the HBA-A mature gene, and the heal site was provided to cleave the DNA fragment at a position of the HPA fragment at a position of the expression of the properties of the prop

Each of the reaction mixtures (50 µl) containing a different 5'-phosphonylated DNA were mixed to make 100 µl, and the mixture was heated in a water bath at 100°C and allowed to cool, to anneal the DNAs. To improve the efficiency of insertion of the annealed phosphonylated DNA into plasmid pUC19, after cleaving the plasmid pUC19 with EcoRI, phosphate groups present at the 5'-ends of the cleaved DNA strands ware eliminated to prevent a resploning of the cleaved plasmid from rejoining during ligation. Namely, 1 µg of pUC19 was treated with 8 units of EcoRII (Nippon gene) at 37°C for 50 minutes in 20 µl of a solution containing 50 mM NaCl, 100 MM Tris-HOL (pH 75) and 7 mM MgCls, to obtain a linearized vector DNA. The reaction mixture was heated at 90°C for 5 minutes to inactivate the enzyme, and to the mixture, 38 µl of water and 1 unit of bacterial alkaline phosphatase was added to make a total volume 60 µl. The reaction mixture was inclusted at 37°C for 50 minutes, the mixture was extracted with phenol, and the resulting aqueous phase was treated with ethanol to precipitate DNA, which was then lyophilized to be used in the next step.

The dephosphorylated linearized vector pUC19 (30 ng) thus prepared and 10 ng of the phosphorylated double-stranded DNA prepared as described above were treated with 28 units of T4 DNA ligasa (Takasa Shuzo) at 15°C for 4 hours in 30 µi of a solution containing 65 mM Tris-HCl (pH 7.5), 6.5 mM MgCls, 10 mM

dithlothreitol and 1 mM ATP to obtain a plasmid.

Competent E. coli cells to be transformed were prepared by a calcium phosphate method of Mandel, M. and Higa, A., J. Mol. Biol. 53, 159-162 (1970). Namely, E. coli TB-1 was cultured overnight in LB medium containing 10 g tryptone, 5 g yeast extract, and 10 g NaCl in 1 f of water (pH 7.4), the culture was diluted 100-fold with the same medium, and culturing was carried out at 37°C with shaking until an ODeoo value reached 0.6. The culture (1.5 ml) was centrifuged at 5000 rpm for 5 minutes to collect cells, the cells were then suspended in 750 µl of 50 mM CaCl2, and after resting on ice for 20 minutes, the suspension was centrifuged to collect the cells. The resulting pellet was resuspended in 100 µl of 50 mM CaCl2, and the above-mentioned DNA ligase reaction mixture was added to the suspension, and the mixture was maintained on ice for 40 minutes. After incubation at 42°C for one minute, 1 ml of LB medium was added to the mixture, which was then incubated at 37°C for 30 minutes. The incubated suspension (0.1 ml) was spread on an X-Gal agar medium prepared by dissolving 155 mg of 5-bromo-4-chloro-3-indolyl-β-galactoside, 10 g of tryptone, 8 g of NaCl and 12 g of agar in 1 ℓ of water and adjusting the pH to 7.2, and incubated overnight at 37°C. Among colonles formed on the agar plate, white colonies were selected, transferred to a fresh agar medium, and cultured overnight. Cells on the agar plate were picked up and inoculated to LB liquid medium and cultured overnight. The culture (1.5 ml) was centrifuged to collect cells. The cells were subjected to mini-preparation of plasmid DNA by a conventional procedure described by Maniatis et al., Molecular Cloning: A laboratory Manual 1982. The resulting plasmid DNA was cleaved with appropriate restriction enzymes, for example, those which cleave restriction sites in the inserted synthetic DNA such as EcoRI, Nael, Hpall, etc., or those which cleave restriction sites in the vector pUC19 such as Pvul, Bgll, Sspl etc., and the cleavage products were analyzed by agarose gel electrophoresis or polyacrylamide gel electrophoresis to determine a size of the inserted DNA. In this manner, a recombinant plasmid which contained a DNA insert having an appropriate size was identified. A DNA fragment containing this DNA insert was introduced again into M13mp phage DNA, and the nucleotide sequence thereof was determined by a dideoxy chain termination method of Sanger, F., Nicklen, S., and Corlson, A.R., Proc., Natl. Acad. Sci. U.S.A. 74, 5463-5467 (1977). A desired plasmid pUS-phoA was identified.

Example 3. Construction of plasmid pUC-phoA-HSA-A (Figs. 2-1 to 2-2)

Piasmid puC-phoA-HSA-A containing DNA coding for a fused protein comprising a signal peptide of E. coli alkaline phosphatase and human normal serum albumin A was constructed as follows.

A clone Agi11 (HSA-II) containing HSA cDNA derived from a human liver cDNA library was cleaved with EcoRI and XbaI to obtain a DNA fragment containing the cDNA. Plasmid pUCI9 was cleaved with EcoRI and XbaI to obtain a larger DNA fragment. These DNA fragments were ligated together using T4 DNA ligase to construct a recombinant plasmid pUCH49A-EX.

The plasmid pUC-HSA-EX was dipested with Ahall and Sall to obtain a smaller DNA fragment which encodes an amino acid sequence from 12h Lys to 558th The 70 human insutre normal serum albumin A. To construct a gene coding for human mature normal serum albumin A, a DNA fragment corresponding to the 5-portion of the mature albumin gene was prepared by ameasing two chemically synthesized oligonucleotides. This DNA fragment has, at the 5'-terminal side thereof, an Hpall cleavage site and Clal cleavage site to provide a cohesive end which can fuse with DNA coding for a signal peptide of alakine phosphatase, and comprises codons coding for an amino acid sequence from the first Asp to 11th Phe. The annealed DNA fragment was phosphorylated at the 5'-end thereof using 14 ponucleotide kinase. On the other hand, a typical E. coli muticioning vector pAT 153 (Amersham; Twigg, AJ. and Sherratt, D, Nature, 283 216-218, 1980) was closeved with Call and Sall, to obtain a larger DNA fragment. The above-prepared three DNA fragments were ligated using 74 DNA ligase to construct a recombinant plasmid pAT-HSA-CX. In this plasmid, DNA coding for the first Asp to 11th Phe 1s tused with DNA coding for for the 12th, by co 556th Phe. The plasmid pAT-HSA-CX was digested with EoRI and Xbal to obtain a smaller DNA fragment to end to 556th Phe. The plasmid pAT-HSA-CX was digested with EoRI and Xbal to obtain a smaller DNA fragment coding for the first Asp to 356th Phe of the human normal serum albumin.

On the other hand, the phage Agt11 (HSAI-A) selected from the human liver cDNA bizary, as described above, was digested with EcoR1 to obtain a DNA fragment containing a cDNA coding for the C-terminal half of the human normal serum albumin A. The DNA fragment was inserted to the EcoR1 site of plasmid pUC18 to construct a recombinant plasmid pUC-HSA-1. This plasmid was digested with Xball and Hindlil to obtain a CDNA fragment containing the region coding for SSSII Leu to the SSSII cauto toys terminal Leu and 3'-terminal non-coding region consisting of 82 nucleotides. On the other hand, a plasmid pUC18 was digested with EcoR1 and Hindlil to obtain a larger fragment. The above-prepared three DNA fragments were ligated using 14 DNA ligase to construct a recombinant plasmid pUC-HSA-CH containing an entire cDNA coding for human mature normal serum albumin.

A nucleotide sequence of cDNA coding for an entire amino acid sequence of human mature normal serum albumin A and a corresponding amino acid sequence are shown in Figs. 3-1 to 3-5.

To join the cDNA coding for human mature normal serum albumin A with DNA coding for a signal peptide of alkaline phosphatase (phoA), a plasmid pUC-HSA-CH was digested with EcoRI and Cial to obtain a larger DNA fragment. A plasmid pUC-phoA was digested with EcoRI and Mspl (recognizing the same sequence as that of Hpal) to obtain a smaller DNA fragment. These DNA fragments were ligated using 14 DNA ligase to construct plasmid pUC-phoA-HSA-A (Fig. 3), which contain DNA coding for a phoA signal peptide consisting of 21 amino acids fused to human mature normal serum albumin A. This plasmid was used to transform E. coll H3101.

Example 4. Construction of expression plasmid pAT-phoA-HSA-A

A plasmid pAT-phoA-HSA-A for expression of human normal serum albumin A was constructed as follows. To express the above-mentioned gene in E. coll, the gene should be linked with an SD sequence responsible for an effective initiation of a translation and a promoter responsible for an effective initiation of a translation. In this example, a trp promoter and trpL SD sequence were used. A vector containing the trp promoter and trpL SD sequence is exemplified by plasmid ph-TNF (Ikehara et al., Chem. Pharm. Bulletin, in press) wherein the trp promoter and trpL SD sequence have been inserted in pBR322. However, to increase the copy number of a recombinant plasmid leading to a gene dosage effect, a plasmid based on plasmid pAT153 (Amersham; Twigg, A.J. and Sherratt, D., Nature, 283, 216-218, 1980) wherein replication poison sequence of pBR322 has been deleted, is preferably used. To this end, a plasmid pheTNF was digested with Pstl and Clal to obtain a DNA fragment containing a trp promoter and trpL SD sequence. On the other hand, a plasmid pAT153 was digested with Pstl and Clal to obtain a larger DNA fragment. Next, these DNA fragments were ligated to construct a plasmid pAT-trp. The plasmid pAT-trp was cleaved at a unique Clai site present downstream of the SD sequence, and resulting cohesive ends were filled in using E. Coll DNA polymerase I, and a resulting linearized plasmid was digested with Sall to obtain a larger fragment.

On the other hand, plasmid pUC-phoA-HSA-A was digested with EcoRI and HindIII to obtain a smaller DNA fragment containing phoA-HSA-A cDNA, which was then ligated to a larger EcoRI/HindIII double digest of pAT 153 to construct a recombinant plasmid pAT-phoA-HSA. This plasmid was digested with EcoRI, and a resulting linearized plasmid was treated with E. coli DNA polymerase I to fill in the ends thereof, and cleaved with Sall to obtain a smaller DNA fragment containing phoA-HSA-A cDNA. This fragment was ligated with the DNA fragment prepared from plasmid PAT-trp, as described above, to construct a recombinant plasmid pAT-trp-phoA-HSA-A. This recombinant plasmid was used to transform E. coli HB101 and E. coli C600 to obtain E. coli HB101 (pAT-trp-phoA-HSA-A) and E. coll C600 (pAT-trp-phoA-HSA-A), respectively. E. coli C600 (pAT-trp-phoA-HSA-A) was deposited with the Fermentation Research Institute Agency of Industrial Science and Technology, 1-3, Higashi 1-chome, Tsukuba-shi, Ibaraki, Japan, on February 17, 1988 as FERM P-9874, and transferred to the international deposited under the Budapest Treaty as FERM BP- 2290 on February 17th, 1989.

Example 5. Production of fused protein

Fused protein comprising a signal peptide of E. coli alkaline phosphatase and human normal serum albumin A was produced using E. coli containing pAT-trp-phoA-HSA-A, as follows.

E. coli C600m r transformed with pAT-trp-phoA-HSA-A was inoculated to 5 ml of Luna broth (Bacto tryptone 1%, yeast extract 0.5%, NaCl 0.5%) supplemented with 25 µl ampicillin, and cultured for 18 hours at 37° C. A part of this culture (0.2 ml) was inoculated to 5 ml of M9-CA medium (Na₂HPO₄ 0.6%, KH₂PO₄ 0.3%, NaCl 0.5%, NH₄Cl 0.1%, CaCl₂ 0.1 mM, MgSO₄ 2 mM, and casamino acid 0.8%) supplemented with 25 μg/ml ampicillin, and culturing was carried out at 37°C for 30 minutes. To the culture was added 20 μg/ml inducer 3-indole acrylic acid (IAA), and culturing was carried out at 37°C for an additional 5 to 7 hours.

Preparation of Insoluble fraction

The culture prepared as described above was centrifuged at 7000 rpm for 5 minutes to collect cells. The precipitated cells were resuspended in 20% sucrose, 25 mM Tris-HCl (pH 7.5), 10 mM EDTA, 1 mM phenylmethanesulfonyl fluoride, and to the suspension was added egg white lysozyme to 0.2 mg/ml. The mixture was allowed to stand at 37°C for 15 minutes to digest the outer membrane, to obtain spheroplasts. The suspension was then cooled in ice, and centrifuged at 10000 rpm for 10 minutes to precipitate the spheroplasts. The spheroplasts were resuspended in a sucrose solution and disrupted in a Polytron homogenizer (dial: 8) in an ice bath. The homogenate was centrifuged at 15,000 rpm for 20 minutes at 4°C to obtain cell debris. The cell debris was resuspended in 25 mM Tris-HCL (pH 7.5), and the suspension was centrifuged at 15,000 rpm for 20 minutes. This operation was repeated once more to obtain a desired insoluble fraction.

SDS-polyacrylamide gel electrophoresis

1) Analysis of whole cellular protein

A part of the culture (0.5 ml) was centrifuged at 7000 rpm for 5 minutes to collect cells. The cells were suspended in 10 µl of SDS-sample solution (62.5 mM Tris-Hcl, pH 6.8, 2% SDS, 10% sucrose, 5% 2-mercaptoethanol, and the suspension was heated at 100°C for 5 minutes. This was subjected to electrophoresis on SDS-polyacrylamide gel (gel concentration, 10%) by a method of Laemmli, Nature (London), 227, 680-685 (1970).

2) Analysis of insoluble fraction

A portion of the insoluble fraction prepared as described above was diluted with the SDS-sample solution, and the suspension was heated at 100°C for 5 minutes to dissolve the insoluble proteins, and subjected to

SDS-acrylamide gel electrophoresis.

3) Staining and destaining

After electrophoresis, the gel was dipped in a staining solution containing 0.25% Coomassie Brilliant Blue, 45% ethanol and 10% acetic acid for 30 to 60 minutes, and then in a destaining solution containing 5% methanol and 10% acetic acid in a destaining apparatus (BioRad, Model 555 type).

Western blottling and immunological detection

After finishing the SDS-PAGE, the get was removed from the glass plate, and a nitrocellulose filter (Bio-Rad, Trans-blot ®) and two 3 MM filter papers (Whatman) were impregnated with a blotting solution (0.3% Tris, 1.44% glycine and 20% methanol). On a pad previously impregnated with the blotting solution, the above-mentioned filter paper, gel, nitrocellulose filter, and filter paper were piled in this order, the upper filter paper was covered with the pad, and the whole was put in a blotting apparatus (TEFCO; Model: TC-608). The apparatus was filled with the blotting solution, and an electrophoresis was carried out at 200 mA for one hour.

After finishing the electrophoresis, the nitrocellulose filter was peeled from the gel and treated in a TBS solution (25 mM Tris-HCl, pH 7.5, 0.5 M NaCl) for 10 minutes. The filter was then treated in a TBS solution containing 3% gelatin for 30 minutes, followed by treatment in TBS containing 0.025% Tween 20 (TTBS solution) for 5 minutes. This procedure was repeated. An IgG fraction of rabbit anti-human albumin serum (Cappel) was diluted 2000-fold with TTBS containing 1% gelatin, and the filter was dipped in this solution for 2 to 18 hours. The sheet was then transferred in TTBS and maintained therein for 5 minutes. This procedure was repeated twice. A horseradish peroxidase-labeled goat anti-rabbit IgG antibody (Bio-Rad) was diluted 3000-fold with TTBS containing 1% gelatin, and the filter was dipped in this solution for 2 hours. Next, the filter was washed twice with TTBS and once with TBS, for 5 minutes each. The filter was dipped in TBS containing 0.015% H₂O₂ , 0.05% HRP color development reagent (Bio-Rad) and 16.7% methanol for 15 minutes, and then dipped in water for 30 minutes. The band of substance which cross-reacted with human normal serum albumin A was colored deep purple on the filter (Fig. 4), and an expression product of the present invention having a molecular weight of 69,000 was detected.

Claims

1 A cDNA coding for human normal serum albumin A having an amino acid sequence represented in Figures 3-1 to 3-5.

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- 2. A cDNA according to claim 1, having a nucleotide sequence represented in Figure 3-1 to 3-5. 3 An expression plasmid comprising a cDNA coding for human normal serum albumin A having an
- amino acid sequence represented in Figures 3-1 to 3-5. 4. An expression plasmid according to claim 4, wherein the cDNA has a nucleotide sequence
- represented in Figures 3-1 to 3-5. 5. A host transformed with an expression plasmid comprising a cDNA coding for human normal serum
- albumin A having an amino acid sequence represented in Figures 3-1 to 3-5. 6. A host according to claim 5, wherein the cDNA has a nucleotide sequence represented in Figures 3-1
- 7. A process for production of human normal serum albumin A comprising the steps of culturing a host transformed with an expression plasmid comprising a cDNA coding for the human normal serum albumin having an amino acid sequence represented in Figures 3-1 to 3-5 to express the protein alone or in a form of a fused protein with another protein, and obtaining the human normal serum albumin A.
- 8. A process according to claim 7, wherein the cDNA has a nucleotide sequence represented in Figure 3-1 to 3-5.



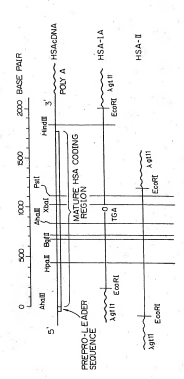
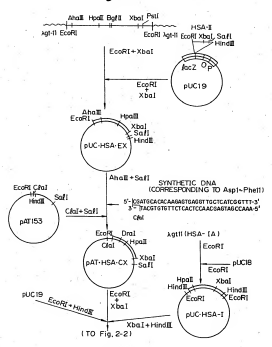
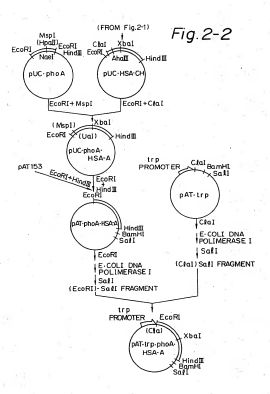


Fig. 2-1





Val GTG	Val	Leu	Ala GCA	Leu TTG	Tyr
Leu TTG	GAA	Thr	Cys	Arg	Lys AAA
Ala GCC	Asn	His	Cys	Pro CCC	Lys
Lys	Val	Leu	Asp	Leu	Leu TTG
Phe	Leu	Ser	Ala GCT	Asn	Phe
Asn	Lys	Lys.	Met	Pro	Thr
Glu	Val GTA	ASP	Glu	Asn	Glu
Glu	His	Cys	Gly	Asp GAC	Glu
GLY	Asp	Asn	TYE	Asp	Asn
Leu TTG	Glu	Glu	Thr	Lys	Asp
Asp	Phe TTT	Ala GCT	Glu	His	His
LYS	Pro	Ser	Arg	Gln	Phe
Phe	Cys	Glu GAG	Leu	Leu	Ala
Arg	Gln	Asp GAT	Thr	Phe	Thr
His	Gln	Ala GCT	Ala GCA	Cys	Cys
Ala GCT	Leu	Val	Val GTT	100 Glu GAA	Met
Val GTT	Tyr	Cys	Thr	Asn AAT	val GTG
Glu GAG	Gln	Thr	Cys TGC	Arg	Asp
Ser	Ala	Lys	Leu	Glu GAG	Val GTT
Lys	Phe	50 Ala GCA	Lys	Pro	Glu
His	Ala GCC	Phe TTT	Asp	Glu	Pro
Ala GCA	Ile	Glu	GLY	Gln	Arg
Asp	Leu	Thr	Phe	Lys	Val

Tyr	GAA	GLY	Glu GAA	Cys	Lys
Arg	Asp	Phe TTT	Ala GCA	Glu	Leu
Lys	Leu	Lys	Phe	CTT	Lys
Ala	Lys	Gln	Glu	250 Leu CTG	Ser
Phe	Pro	CTC	Ala GCT	Asp	Ser
Phe	Leu TTG	Ser	Lys	GLY	ATC
Leu	Leu	Ala GCC	Pro	His	Ser
Leu	Cys	Cys TGT	Phe	Cys	Asp
Glu	Ala GCC	Lys	Arg	Cys	Gln
Pro	Ala	Leu	Gln	Glu	Asn
Ala	Lys	Arg	Ser	Thr	Glu
150 Tyr TAT	Asp	Gln	Leu	His	Cys
Phe TTT	Ala GCT	Lys	Arg	Val	Ile
Tyr	Ala	Ala GCC	Ala GCT	LYS	Tyr
Pro	Gln	Ser	Val GTA	Thr	Lys
His	Cys	Ser	Ala	Leu	Ala
Arg	Cys	Ala	Trp	Asp	Leu
Arg	Glu	Lys	Ala GCA	Thr	Asp
Ala GCC	Thr	G1y GGG	Lys	Val GTG	Ala GCG
Ile	Phe TTT	Glu	Phe	Leu TTA	Arg
Glu	Ala GCT	Asp	Ala GCT	LYS	Asp
TYT	Ala	Arg	Arg	Ser	Asp
Leu	Lys	Leu	Glu	Val GTT	Ala GCT

Pro	Ala GCA	Leu	Glu	Asn	Lys
Met	Glu	Val	His	Gln	Lys
Glu	Ala GCT	Val GTC	Pro	Lys	Thr
Asp	TYr	Ser	Asp GAT	Ile	Tyr
Asn	Asn	Tyr	Ala GCA	Leu	Arg
Glu	Lys	Asp	Ala	Asn	Val GTT
Val GTG	Cys	Pro	Ala GCC	Gln	Leu
Glu	Val	His	Cys	Pro	Leu
Ala	Asp	Arg	Cys	GAG	Ala
Ile	Lys	Arg	Lys	Glu	Asn
Cys	Ser	Ala GCA	Glu	val GTG	Gln
His	Glu	TYT	Leu	Leu	Phe
Ser	Val	GLu	Thr	Pro	Lys
Lys	Phe	Tyr	Thr	Lys	Tyr
Glu	Asp	Leu	Glu	Phe	400 Glu GAG
Leu	Ala	Phe	TYL	Glu	Gly
Leu	Ala	Met	Thr	Asp	Leu
Pro	Leu	GLY	Lys	Phe	Gln
Lys	Ser	Leu	350 Ala GCC	Val	Glu
Glu	Pro	Phe	Leu	LYS	Phe
Cys	Leu	Val	Arg	Ala GCC	Leu
Cys	Asp	Asp GAT	Leu	Tyr	610
Glu	300 Ala GCT	Lys	Leu	Cys	Cys

Leu Asn Thr Leu Ala Lys Gln Val Glu Ala Phe Ser Asn Leu Ala Thr Asp GLY Leu Ser Asn Gln Asp Val Val Glu Phe Lys Met Val Thr Glu Lys Val Gly Ser Cys 500 Lys Ile Ala Leu Leu Cys Pro Gln Lys Asn TYT Lys Val Arg Leu Arg Asp Tyr Glu Thr Gln Ser 450 Glu GAA Val Thr Lys Glu Val Ala GCA Arg Glu Glu Lys Glu Asp Cys Asp Ser Thr Val Pro Ser Val Leu Ala GCA Leu Met Glu Val Thr Lys Thr Arg Pro Leu Cys Pro Pro Lys Thr Ala GCT Ile Lys Thr Ala GCA Lys Ser Asp His Ser Glu Glu Phe Ala Lys Val Pro His Cys His Val Gln His Leu Pro Phe Leu Pro Lys Val Arg Thr Glu Val Cys Cys Arg Phe Val

Ala Glu Glu Gly Lys Lys Leu GCC GAG GAG GGT AAA AAA CTT Ala Phe Val Glu Lys Cys Cys Lys Ala Asp Asp Lys Glu Thr Cys Phe GCT TTT GTA GAG AAG TGC TGC AAG GCT GAC GAT AAG GAG ACC TGT TTT Ala Ser Gln Ala Ala Leu Gly Leu End GCA AGT CAA GCT GCC TTA GGC TTA TAA Val Ala A

Fig. 4

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Fig. 5

HSA -1 5-AAGGAAATAAAGGTTACCCACTTCATTGTGCCAAAGGC-3 REGION CORRESPONDING TO 5-NON-CODING REGION-MEII-LEU9 (12 NUCLEOTIDES)

HSA-2 5'-AAGGTCCGCCCTGTCACCACCATTCAAGCAGATCTCC-3' REGION CORRESPONDING TO 61 y 248~Leu 260

REGION CORRESPONDING TO VA! 576~Leu 585~3' NON-CODING REGION (6 NUCLEOTIDES) HSA-35'-TAGATGTTATAAGCCTAAGGCAGCTTGACTTGCAGCAAC-3'

(1) Publication number:

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- (2) Representative: Bizley, Richard Edward et al BOULT, WADE & TENNANT 27 Furnival Street London EC4A 1PO(GB)
- A cDNA coding for human normal serum albumin a, and a process for production of the albumin.
- A cDNA coding for human normal serum abumin A; an expression vector comprising the cDNA coding for human normal serum abumin A; a host transformed with the expression vector comprising the cDNA coding for human normal serum albumin A; and a process for the production of the human normal serum albumin A comprising the steps of culturing a host transformed with an expression vector comprising a cDNA coding for the human normal serum albumin to express the protein alone or in a form of a fused protein with another protein, and obtaining the human normal serum albumin A.

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ROPEAN SEARCH REPORT

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	Citation of document with indic	ERED TO BE RELEV	Relevant	CLASSIFICATION OF THE
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X	EP-A-0 200 590 (GENE	TICA)	1-8	C 12 N 1/20
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^.	* Claims *	0.00111.0017		-
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